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L2 1 SEA FILE=HCAPLUS ABB=ON PLU=ON L1
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L2 (ANSWER 1 OF 1 | HCAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

2004:267260 HCAPLUS

DOCUMENT NUMBER:

140:297533

TITLE:

Peptides and related molecules that modulate nerve

, growth factor activity

INVENTOR(S):

Boone, Thomas C.; Wild, Kenneth D., Jr.; Sitney, Karen

C.; Min, Hosung; Kimmel, Bruce

PATENT ASSIGNEE(S):

Amgen Inc., USA

SOURCE:

PCT Int. Appl., 267 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA.	PATENT NO. WO 2004026329						APPLICATION NO.					DATE					
WO							WO 2003-US29866					20030919					
	W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
		CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
		PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	TJ,	TM,	TN,	TR,	TT,	TZ,
		UA,	UG,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW							
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US	US 2004121959				A1	;	2004	0624	1	US 2	003-	6664	80		2	0030	918
PRIORIT'	Y APP	LN.	INFO	.:					1	US 2	002-	4125	24P		P 2	0020	919
									1	US 2	003-	6664	80	i	A 2	0030	918

OTHER SOURCE(S): MARPAT 140:297533

- AB The present invention relates to certain biol. active peptides and polypeptides which can be used as therapeutics or prophylactics against diseases or disorders linked to nerve growth factor (NGF) as the causative agent. In one aspect of the present invention, pharmacol. active polypeptides comprising peptides linked to one or more Fc domains are provided.
- IC ICM A61K038-10
- ICS A61K038-16; C07H021-04; C07K007-08; C07K014-00
- CC 1-11 (Pharmacology)
- TT 57-88-5D, Cholesterol, conjugates with peptides 7093-67-6D, conjugates with peptides and Fc domains 18861-82-0D, conjugates with peptides and Fc domains 25322-68-3D, Polyethylene glycol, conjugates with peptides 676329-46-7D, linker-peptide-Fc domain conjugates 676329-48-9D, linker-peptide-Fc domain conjugates 676329-50-3D, linker-peptide-Fc domain conjugates 676329-53-6D, linker-peptide-Fc domain conjugates 676329-54-7D, linker-peptide-Fc domain conjugates 676329-55-8D, linker-peptide-Fc



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                                 676373-34-5
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     (Biological study); USES (Uses)
        (peptides and related mols. that modulate nerve growth factor activity
        linked to vehicles such as antibody Fc domains for treatment of diseases associated with pain)
REFERENCE COUNT:
                                THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS
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RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT



Russel 10/666,480

01/07/2005

William

WHI I

WHAT

Searched by Paul Schulwitz 571-272-2527

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OM protein - protein search, using sw model

Run on: January 7, 2005, 15:34:01; Search time 62 Seconds

(without alignments)

167.044 Million cell updates/sec

Title: US-10-666-480-231

Perfect score: 104

Sequence: 1 DQLGDWMLNYFRLVPPGT 18

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 1825181 seqs, 575374646 residues

Total number of hits satisfying chosen parameters: 1825181

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database: UniProt 02:*

1: uniprot_sprot:*
2: uniprot_trembl:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

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No.	Score	Match	Length	DB	ID	Description
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3	50	48.1	343	2	BAC83487	Bac83487 oryza sat
4	50	48.1	518	2	Q9VLU7	Q9vlu7 drosophila
5	50	48.1	552	2	Q96VC4	Q96vc4 emericella
6	50	48.1	762	2	Q6R7C7	Q6r7c7 ostreid her
7	50	48.1	762	2	AAS00988	Aas00988 ostreid h
8	48	46.2	243	2	084617	084617 chlamydia t
9	48	46.2	297	2	Q82MC4	Q82mc4 streptomyce
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21	46	44.2	362	2	Q8FND8	Q8fnd8 corynebacte
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23	46	44.2	417	1	RNBP_HUMAN	P51606 homo sapien
24	46	44.2	419	1	RNBP MOUSE	P82343 mus musculu
25	46	44.2	464	2	Q9KEM8	
26	46	44.2	554	2	Q7RXW8	Q9kem8 bacillus ha
27	46	44.2	1394	2	Q7Q971	Q7rxw8 neurospora
28	46	44.2	2042	2	Q7Q3V4	Q7q971 anopheles g
20 29	45.5	44.2			- ·-	Q7q3u4 anopheles g
			183	2	Q89K86	Q89k86 bradyrhizob
30	45.5	43.8	1202	2	Q7WF28	Q7wf28 bordetella
31	45.5	43.8	1224	2	Q7W3Q1	Q7w3q1 bordetella
32	45	43.3	142	2	Q9S2F7	Q9s2f7 streptomyce
33	45	43.3	331	2	Q6C1P0	Q6c1p0 yarrowia li
34	45	43.3	359	2	Q8RIS8	Q8ris8 actinoplane
35	45	43.3	453	2	Q8EKN5	Q8ekn5 shewanella
36	45	43.3	878	1	ACON_RICCN	Q92g90 rickettsia
37	45	43.3	878	1	ACON_RICPR	Q9zcf4 rickettsia
38	45	43.3	878	2	Q7P9X6	Q7p9x6 rickettsia
39	45	43.3	890	2	Q83B05	Q83b05 coxiella bu
40	45	43.3	928	2	Q6FUJ3	Q6fuj3 candida gla
41	44.5	42.8	100	2	Q7UZC7	Q7uzc7 rhodopirell
42	44.5	42.8	324	2	Q876L8	Q87618 trichoderma
43	44.5	42.8	421	2	Q9GRS9	Q9grs9 leishmania
44	44.5	42.8	486	2	Q00995	Q00995 kluyveromyc
45	44.5	42.8	486	2	Q6CWY3	Q6cwy3 kluyveromyc

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RESULT 1
Q9PJC8
ID
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                                    PRT;
                                           243 AA.
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АC
DT
     01-OCT-2000 (TrEMBLrel. 15, Created)
\mathsf{DT}
     01-OCT-2000 (TrEMBLrel. 15, Last sequence update)
DT
     01-MAR-2004 (TrEMBLrel. 26, Last annotation update)
DE
     Hypothetical protein TC0901.
     OrderedLocusNames=TC0901;
GN
OS
     Chlamydia muridarum.
OC
     Bacteria; Chlamydiae; Chlamydiales; Chlamydiaceae; Chlamydia.
OX
     NCBI TaxID=83560;
RN
RP
     SEQUENCE FROM N.A.
RC
     STRAIN=MoPn / Nigg;
RX
     MEDLINE=20150255; PubMed=10684935;
RA
     Read T.D., Brunham R.C., Shen C., Gill S.R., Heidelberg J.F.,
RA
     White O., Hickey E.K., Peterson J.D., Utterback T.R., Berry K.J.,
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     Bass S., Linher K.D., Weidman J.F., Khouri H.M., Craven B., Bowman C.,
RA
     Dodson R.J., Gwinn M.L., Nelson W.C., DeBoy R.T., Kolonay J.F.,
RA
     McClarty G., Salzberg S.L., Eisen J.A., Fraser C.M.;
RT
     "Genome sequences of Chlamydia trachomatis MoPn and Chlamydia
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RT
    pneumoniae AR39.";
RL
    Nucleic Acids Res. 28:1397-1406(2000).
    EMBL; AE002356; AAF39694.1; -.
DR
DR
    PIR; E81652; E81652.
    TIGR; TC0901; -.
DR
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DR
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    Pfam; PF01996; DUF129; 1.
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KW
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                         50.0%; Score 52; DB 2; Length 243;
 Query Match
 Best Local Similarity 53.8%; Pred. No. 6.9;
 Matches
          7; Conservative 5; Mismatches 1; Indels
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           1 DQLGDWMLNYFRL 13
Qу
             :1:1:1: | | | | | |
Db
         110 NQIGEWLKNYFRV 122
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Search completed: January 7, 2005, 16:05:32 Job time: 65 secs

OM protein - protein search, using sw model

January 7, 2005, 15:52:11; Search time 16 Seconds Run on:

(without alignments)

108.244 Million cell updates/sec

Title: US-10-666-480-231

Perfect score: 104

Sequence: 1 DQLGDWMLNYFRLVPPGT 18

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

283416 seqs, 96216763 residues Searched:

Total number of hits satisfying chosen parameters: 283416

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : PIR 79:*

1: pir1:*

2: pir2:*

3: pir3:*

4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

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	No.	Score	Match	Length	DB	ID	Description
	1	52	50.0	243	2	E81652	conserved hypothet
	2	50	48.1	552	2	JC7666	serine-type carbox
	3	48	46.2	243	1	G71493	hypothetical prote
	4	47	45.2	405	2	T44249	transport protein
	5	47	45.2	419	2	JX0187	renin-binding prot
	6	47	45.2	903	2	A87704	aconitate hydratas
	7	46	44.2	315	2	F84683	hypothetical prote
	8	46	44.2	417	2	JX0188	renin-binding prot
	9	46	44.2	464	2	Н83752	glutamate dehydrog
	10	45	43.3	142	2	T36147	probable regulator
	11	45	43.3	878	2	A97854	aconitate hydratas
	.12	45	43.3	878	2	A71641	aconitate hydratas
	13	44.5	42.8	486	2	S72516	FOG1 protein - yea

14	44	42.3	944	2	G87135	aconitate hydratas
15	44	42.3	1272	2	Т37807	hypothetical prote
16	43	41.3	137	2	G95352	protein [imported
17	43	41.3	173	2	C86503	acyltransferase [i
18	43	41.3	246	2	A72040	ct611 hypothetical
19	43	41.3	246	2	F86585	CT611 hypothetical
20	43	41.3	246	2	A81502	conserved hypothet
21	43	41.3	273	2	s75316	phycocyanin-associ
22	43	41.3	507	2	T31975	hypothetical prote
23	43	41.3	1270	2	T09194	adaptor protein in
24	42.5	40.9	219	2	B75533	hypothetical prote
25	42.5	40.9	402	2	AG0638	probable membrane
26	42.5	40.9	412	2	F64849	probable permease
27	42.5	40.9	412	2	C99809	hypothetical prote
28	42.5	40.9	668	2	AG1989	hypothetical prote
29	42.5	40.9	780	2	G72724	probable penicilli
30	42	40.4	411	2	B71451	probable desucciny
31	42	40.4	418	2	B72245	conserved hypothet
32	42	40.4	420	2	AB0246	glutamyl-tRNA redu
33	42	40.4	552	2	G69464	acetolactate synth
34	42	40.4	742	2	H95270	probable kinase/es
35	42	40.4	763	2	A29562	PRT1 protein - yea
36	42	40.4	810	2	T33323	hypothetical prote
37	42	40.4	890	2	AH0270	aconitate hydratas
38	42	40.4	895	2	AI3483	aconitate hydratas
39	42	40.4	897	2	AD2906	aconitate hydratas
40	42	40.4	897	2	E97681	aconitate hydratas
41	41.5	39.9	412	2	G85668	hypothetical prote
42	41.5	39.9	796	2	D97065	transketolase [imp
43	41	39.4	165	2	T46052	ADP-ribosylation f
44	41	39.4	321	2	AH2766	conserved hypothet
45	41	39.4	321	2	C97547	hypothetical prote

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RESULT 1
E81652
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conserved hypothetical protein TC0901 [imported] - Chlamydia muridarum (strain

C; Species: Chlamydia muridarum, Chlamydia trachomatis MoPn

C;Date: 31-Mar-2000 #sequence revision 31-Mar-2000 #text_change 09-Dec-2002

C; Accession: E81652

R; Read, T.D.; Brunham, R.C.; Shen, C.; Gill, S.R.; Heidelberg, J.F.; White, O.; Hickey, E.K.; Peterson, J.; Utterback, T.; Berry, K.; Bass, S.; Linher, K.; Weidman, J.; Khouri, H.; Craven, B.; Bowman, C.; Dodson, R.; Gwinn, M.; Nelson, W.; DeBoy, R.; Kolonay, J.; McClarty, G.; Salzberg, S.L.; Eisen, J.; Fraser, C.M.

Nucleic Acids Res. 28, 1397-1406, 2000

A; Title: Genome sequences of Chlamydia trachomatis MoPn and Chlamydia pneumoniae AR39.

A; Reference number: A81500; MUID: 20150255; PMID: 10684935

A; Accession: E81652 A; Status: preliminary A; Molecule type: DNA A; Residues: 1-243 <TET>

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A;Cross-references: GB:AE002356; GB:AE002160; NID:g7190921; PIDN:AAF39694.1;
PID:g7190928; GSPDB:GN00121; TIGR:TC0901
A; Experimental source: strain Nigg (MoPn)
C; Genetics:
A; Gene: TC0901
C; Superfamily: uncharacterized conserved protein
                         50.0%; Score 52; DB 2; Length 243;
 Query Match
 Best Local Similarity 53.8%; Pred. No. 0.94;
           7; Conservative 5; Mismatches 1; Indels 0; Gaps
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Qу
             :1:1:1: | | | | | |
Db
         110 NQIGEWLKNYFRV 122
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Search completed: January 7, 2005, 16:08:34 Job time: 18 secs

3

OM protein - protein search, using sw model

January 7, 2005, 16:08:22; Search time 145 Seconds Run on:

(without alignments)

44.761 Million cell updates/sec

Title: US-10-666-480-231

Perfect score: 104

Sequence: 1 DQLGDWMLNYFRLVPPGT 18

Scoring table: BLOSUM62

Gapop 10.0, Gapext 0.5

Searched: 1603904 seqs, 360571292 residues

Total number of hits satisfying chosen parameters: 1603904

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Published Applications AA:*

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/cgn2 6/ptodata/2/pubpaa/PCTUS PUBCOMB.pep:*

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11: /cgn2 6/ptodata/2/pubpaa/US09C PUBCOMB.pep:*

/cgn2 6/ptodata/2/pubpaa/US09 NEW PUB.pep:* 12:

13: /cgn2_6/ptodata/2/pubpaa/US10A PUBCOMB.pep:*

14: /cgn2 6/ptodata/2/pubpaa/US10B PUBCOMB.pep:*

15: /cgn2_6/ptodata/2/pubpaa/US10C_PUBCOMB.pep:*

16: /cgn2_6/ptodata/2/pubpaa/US10D_PUBCOMB.pep:*

17: /cgn2 6/ptodata/2/pubpaa/US10 NEW PUB.pep:*

18: /cgn2 6/ptodata/2/pubpaa/US11 NEW PUB.pep:*

19: /cgn2 6/ptodata/2/pubpaa/US60 NEW PUB.pep:*

/cgn2_6/ptodata/2/pubpaa/US60 PUBCOMB.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	Query Match	Length	DB	ID	Description
1	104	100.0	18	16	US-10-666-480-231	Sequence 231, App
2	104	100.0	20	16	US-10-666-480-221	Sequence 221, App
3	104	100.0	22	16	US-10-666-480-202	Sequence 202, App
4	104	100.0	22	16	US-10-666-480-237	Sequence 237, App
5	104	100.0	23	16	US-10-666-480-211	Sequence 211, App
6	104	100.0	23	16	US-10-666-480-272	Sequence 272, App
7	104	100.0	24	16	US-10-666-480-219	Sequence 219, App
8	99	95.2	19	16	US-10-666-480-227	Sequence 227, App
9	54	51.9	26	16	US-10-666-480-271	Sequence 271, App
10	51	49.0	67	17	US-10-425-115-316169	Sequence 316169,
11	50	48.1	26	16	US-10-666-480-226	Sequence 226, App
12	50	48.1	343	16	US-10-437-963-193691	Sequence 193691,
13	50	48.1	429	14	US-10-369-493-13233	Sequence 13233, A
14	49	47.1	344	9	US-09-999-672-11	Sequence 11, Appl
15	49	47.1	344	13	US-10-040-863-11	Sequence 11, Appl
16	48	46.2	22	16	US-10-666-480-251	Sequence 251, App
17	48	46.2	207	9	US-09-841-132-300	Sequence 300, App
18	48	46.2	207	17	US-10-872-155-300	Sequence 300, App
19	48	46.2	243	9	US-09-841-132-579	Sequence 579, App
20	48	46.2	243	17	US-10-872-155-579	Sequence 579, App
21	48	46.2	249	9	US-09-841-132-365	Sequence 365, App
22	48	46.2	249	17	US-10-872-155-365	Sequence 365, App
23	48	46.2	297	14	US-10-156-761-9275	Sequence 9275, Ap
24	48	46.2	896	14	US-10-217-096-6	Sequence 6, Appli
25	48	46.2	896	16	US-10-408-765A-620	Sequence 620, App
26	48	46.2	896	17	US-10-757-262-114	Sequence 114, App
27	48	46.2	896	17	US-10-768-158-36	Sequence 36, Appl
28	47	45.2	26	16	US-10-666-480-239	Sequence 239, App
29	47	45.2	179	15	US-10-424-599-162543	Sequence 162543,
30	47	45.2	882	14	US-10-369-493-7994	Sequence 7994, Ap
31	47	45.2	891	14	US-10-369-493-21094	Sequence 21094, A
32	47	45.2	896	14	US-10-369-493-209	Sequence 209, App
33	47	45.2	903	14	US-10-369-493-17080	Sequence 17080, A
34	46	44.2	145	16	US-10-767-701-55257	Sequence 55257, A
35	46	44.2	291	14	US-10-286-264-98	Sequence 98, Appl
36	46	44.2	291	15	US-10-412-699B-916	Sequence 916, App
37	46	44.2	417	14	US-10-116-275-224	Sequence 224, App
38	46	44.2	464	14	US-10-369-493-17155	Sequence 17155, A
39	46	44.2	539	14	US-10-369-493-3911	Sequence 3911, Ap
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41 42	45 45	43.3 43.3	211	17	US-10-425-115-358149	Sequence 358149,
42	44.5	43.3	887 1592	14 16	US-10-369-493-17785	Sequence 17785, A
44	44.5	42.8	1975	16	US-10-437-963-188419 US-10-437-963-188418	Sequence 188419,
45	44.5	42.3	21	16	US-10-666-480-210	Sequence 188418,
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RESULT 1 US-10-666-480-231

[;] Sequence 231, Application US/10666480 ; Publication No. US20040121959A1

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; GENERAL INFORMATION:
; APPLICANT: Boone, Thomas C
 APPLICANT: Wild, Kenneth D
; APPLICANT: Sitney, Karen C
; APPLICANT: Min, Hosung
; APPLICANT: Kimmel, Bruce
; TITLE OF INVENTION: Peptides and Related Molecules That Modulate Nerve Growth
Factor Activity
; FILE REFERENCE: A-827US
; CURRENT APPLICATION NUMBER: US/10/666,480
; CURRENT FILING DATE: 2003-09-18
  PRIOR APPLICATION NUMBER: 60/412,524
  PRIOR FILING DATE: 2002-09-19
; NUMBER OF SEQ ID NOS: 286
 SOFTWARE: PatentIn version 3.1
; SEQ ID NO 231
   LENGTH: 18
   TYPE: PRT
   ORGANISM: Artificial Sequence
   FEATURE:
   OTHER INFORMATION: Therapeutically active peptide of randomly generated,
   OTHER INFORMATION: lly occurring sequence
US-10-666-480-231
  Query Match 100.0%; Score 104; DB 16; Length 18; Best Local Similarity 100.0%; Pred. No. 3.1e-09;
  Matches 18; Conservative 0; Mismatches 0; Indels 0; Gaps
           1 DQLGDWMLNYFRLVPPGT 18
Qу
             Db
           1 DQLGDWMLNYFRLVPPGT 18
RESULT 10
US-10-425-115-316169
; Sequence 316169, Application US/10425115
; Publication No. US20040214272A1
; GENERAL INFORMATION:
; APPLICANT: La Rosa, Thomas J.
; APPLICANT: Kovalic, David K.
; APPLICANT: Zhou, Yihua
; APPLICANT: Cao, Yongwei
; TITLE OF INVENTION: Nucleic Acid Molecules and Other Molecules Associated
With
; TITLE OF INVENTION: Plants
; FILE REFERENCE: 38-21(53222)B
; CURRENT APPLICATION NUMBER: US/10/425,115
; CURRENT FILING DATE: 2003-04-28
; NUMBER OF SEQ ID NOS: 369326
; SEQ ID NO 316169
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  FEATURE:
; NAME/KEY: unsure
  LOCATION: (1)..(67)
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; OTHER INFORMATION: unsure at all Xaa locations
   FEATURE:
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OTHER INFORMATION: Clone ID: MRT4577_51416C.1.pep US-10-425-115-316169

Query Match 49.0%; Score 51; DB 17; Length 67; Best Local Similarity 50.0%; Pred. No. 2.5;

Matches 8; Conservative 4; Mismatches 4; Indels 0; Gaps 0;

3 LGDWMLNYFRLVPPGT 18 Qу :111:11:1 10 MGDWILNHFFFLTPRT 25 Db

Search completed: January 7, 2005, 16:19:54

Job time : 146 secs

OM protein - protein search, using sw model

Run on: January 7, 2005, 15:54:26; Search time 21 Seconds

(without alignments)

56.844 Million cell updates/sec

Title: US-10-666-480-231

Perfect score: 104

Sequence: 1 DQLGDWMLNYFRLVPPGT 18

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 478139 seqs, 66318000 residues

Total number of hits satisfying chosen parameters: 478139

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : Issued Patents AA:*

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4: /cgn2 6/ptodata/1/iaa/6B COMB.pep:*

5: /cgn2_6/ptodata/1/iaa/PCTUS_COMB.pep:*

6: /cgn2 6/ptodata/1/iaa/backfiles1.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

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	4	50	48.1	557	1	US-08-608-267-4	Sequence 4, Appli
	5	50	48.1	557	1	US-08-608-452-2	Sequence 2, Appli
	6	50	48.1	557	1	US-08-608-452-4	Sequence 4, Appli
	7	50	48.1	557	1	US-08-608-224-2	Sequence 2, Appli
	8	50	48.1	557	1	US-08-608-224-4	Sequence 4, Appli
	9	50	48.1	557	2	US-08-967-149-2	Sequence 2, Appli
	10	50	48.1	557	2	US-08-967-149 - 4	Sequence 4, Appli
	11	49	47.1	344	3	US-09-298-886-11	Sequence 11, Appl

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             38.0
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                             UŞ-09-118-442-11
                                                        Sequence 11, Appl
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RESULT 1
US-08-309-341-2
; Sequence 2, Application US/08309341
; Patent No. 5594119
   GENERAL INFORMATION:
    APPLICANT: Yaver, Debbie Sue
    APPLICANT: Thompson, Sheryl Ann
;
    TITLE OF INVENTION: GENE ENCODING CARBOXYPEPTIDASE OF ASPERGILLUS NIGER
    NUMBER OF SEQUENCES: 4
    CORRESPONDENCE ADDRESS:
      ADDRESSEE: No. 55941190 No. 5594119disk of No. 5594119th America, Inc.
       STREET: 405 Lexington Avenue, Suite 6400
;
      CITY: New York
      STATE: New York
      COUNTRY: U.S.A.
           10174-6401
;
    COMPUTER READABLE FORM:
;
      MEDIUM TYPE: Floppy disk
```

```
COMPUTER: IBM PC compatible
;
      OPERATING SYSTEM: PC-DOS/MS-DOS
      SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)
    CURRENT APPLICATION DATA:
     APPLICATION NUMBER: US/08/309,341
      FILING DATE: 16-SEP-1994
      CLASSIFICATION: 435
    ATTORNEY/AGENT INFORMATION:
      NAME: Lowney, Karen A.
      REGISTRATION NUMBER: 31,274
      REFERENCE/DOCKET NUMBER: 4247.000-US
    TELECOMMUNICATION INFORMATION:
      TELEPHONE: 212 867 0123
      TELEFAX: 212 867 0298
  INFORMATION FOR SEQ ID NO: 2:
    SEQUENCE CHARACTERISTICS:
      LENGTH: 557 amino acids
      TYPE: amino acid
      STRANDEDNESS: single
      TOPOLOGY: linear
    MOLECULE TYPE: protein
    ORIGINAL SOURCE:
      ORGANISM: Aspergillus Niger
US-08-309-341-2
 Query Match
                        48.1%; Score 50; DB 1; Length 557;
 Best Local Similarity 75.0%; Pred. No. 7.3;
 Matches 9; Conservative 0; Mismatches 3; Indels 0; Gaps
           4 GDWMLNYFRLVP 15
Qу
             1111 | 1111
Db
        445 GDWMKPYHRLVP 456
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Search completed: January 7, 2005, 16:09:00 Job time: 22 secs

OM protein - protein search, using sw model

Run on: January 7, 2005, 15:49:26; Search time 154 Seconds

(without alignments)

41.929 Million cell updates/sec

Title: US-10-666-480-231

Perfect score: 104

Sequence: 1 DQLGDWMLNYFRLVPPGT 18

Scoring table: BLOSUM62

Gapop 10.0 , Gapext 0.5

Searched: 2002273 seqs, 358729299 residues

Total number of hits satisfying chosen parameters: 2002273

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database: A_Geneseq_23Sep04:*

1: geneseqp1980s:*

2: geneseqp1990s:*

3: geneseqp2000s:*

4: geneseqp2001s:*

5: geneseqp2002s:*

6: geneseqp2003as:*

7: geneseqp2003bs:*

8: geneseqp2004s:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

			8					
ŀ	Result		Query					
	No.	Score	Match	Length	DB	ID	Description	
	1	104	100.0	18	- 8	ADM17879	Adm17879 Nerve gro	,
	2	104	100.0	20	8	ADM17869	Adm17869 Nerve gro	
	3	104	100.0	22	8	ADM17885	Adm17885 Nerve gro	
	4	104	100.0	22	8	ADM17850	Adm17850 Nerve gro	,
	5	104	100.0	23	8	ADM17859	Adm17859 Nerve gro	,
	6	104	100.0	23	8	ADM17920	Adm17920 Nerve gro	,
	7	104	100.0	24	8	ADM17867	Adm17867 Nerve gro	
	8	99	95.2	19	8	ADM17875	Adm17875 Nerve gro	
	9	54	51.9	26	8	ADM17919	Adm17919 Nerve gro	

				_			
10	50	48.1	26	8	ADM17874	Adm17874 Nerve gro	
11	50	48.1	518	4	ABB63107	Abb63107 Drosophil	
12	50	48.1	536	5	ABR38819	Abr38819 A. niger	
13	50	48.1	556	2	AAR96737	Aar96737 A. niger	
14	50	48.1	557	2	AAR96738	Aar96738 A. niger	
15	49	47.1	344	3	AAB36106	Aab36106 Human Sec	
16	49	47.1	344	6	ABG72379	Abg72379 Human Sec	
17	49	47.1	344	7	ADB97786	Adb97786 Human alp	
18	48	46.2	22	8	ADM17899	Adm17899 Nerve gro)
19	48	46.2	164	3	AAB44991	Aab44991 Human sec	:
20	48	46.2	164	3	AAB44992	Aab44992 Human sec	:
21	48	-46.2	198	2	AAY37590	Aay37590 Amino aci	
22	48	46.2	207	3	AAB13696	Aab13696 Chlamydia	į
23	48	46.2	207	4	AAG83264	Aag83264 Protein e	
24	48	46.2	207	5	ABB94235	Abb94235 Chlamydia	Ĺ
25	48	46.2	243	5	ABB94356	Abb94356 Chlamydia	
26	48	46.2	249	5	ABB94257	Abb94257 Chlamydia	
27	48	46.2	470	8	ADH22542	Adh22542 Human tra	
28	48	46.2	707	4	AAM23744	Aam23744 Human EST	
29	48	46.2	707	4	AAM23715	Aam23715 Human EST	ı
30	48	46.2	896	6	ABR43913	Abr43913 Human sod	l
31	48	46.2	896	7	ADJ68814	Adj68814 Human hea	
32	47	45.2	26	8	ADM17887	Adm17887 Nerve gro	
33	47	45.2	419	2	AAR79930	Aar79930 Porcine a	
34	47	45.2	654	8	ADN48015	Adn48015 Thermococ	:
35	47	45.2	892	6	ABM67426	Abm67426 Photorhab	
36	46	44.2	291	7	ADC46699	Adc46699 Thalecres	
37	46	44.2	291	8	ADO02503	Ado02503 Thalecres	
38	46	44.2	417	2	AAR79929	Aar79929 Porcine a	
39	45	43.3	359	5	AAM49555	Aam49555 Actinopla	
40	44	42.3	21	8	ADM17858	Adm17858 Nerve gro	
41	44	42.3	22	8	ADM17880	Adm17880 Nerve gro	
42	44	42.3	687	4	ABB64842	Abb64842 Drosophil	
43	43	41.3	26	8	ADM17861	Adm17861 Nerve gro	
44	43	41.3	164	5	ABP41930	Abp41930 Human ova	
45	43	41.3	246	5	ABB94321	Abb94321 Chlamydia	
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XX
AC
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XX
DT
     17-JUN-2004
                  (first entry)
XX
DE
    Nerve growth factor activity modulating peptide SEQ ID NO:231.
XX
KW
    nerve growth factor modulator; NGF modulator; analgesic; NGF inhibitor;
KW
    nerve growth factor inhibitor; neurologic pain; diabetic neuropathy;
KW
     post-herpetic neuralgia; inflammatory pain; migraine; asthma;
KW
    hyperactive bladder; psoriasis; cancer; acute pain; dental pain;
KW
    surgical pain; pain; causalgia; demyelinating disease;
KW
    trigeminal neuralgia; chronic alcoholism; stroke; thalamic pain syndrome;
```

KW diabetes; acquired immuno deficiency syndrome; AIDS; headache; KW inflammation; arthritis; rheumatic disease; lupus; osteoarthritis; KW inflammatory bowel disorder; inflammatory eye disorder; sunburn; KW carditis; dermatitis; myositis; neuritis; collagen vascular disease; chronic inflammatory condition; neuropathic pain; genitourinary; wound; KW KW burn; allergic skin reaction; pruritus; vitiligo; KW gastrointestinal disorder; colitis; gastric ulceration; duodenal ulcer. XX OS Synthetic. XX PN WO2004026329-A1. XX PD 01-APR-2004. XX PF 19-SEP-2003; 2003WO-US029866. XX 19-SEP-2002; 2002US-0412524P. PR 18-SEP-2003; 2003US-00666480. PR XX PA (AMGE-) AMGEN INC. XX PΙ Boone TC, Wild KD, Sitney KC, Min H, Kimmel B; XX DR WPI; 2004-283150/26. XX PT

Novel peptide capable of modulating nerve growth factor activity, useful for treating disease or disorder e.g., acute pain, dental pain, cancer, migraine and collagen vascular disease.

Claim 113; SEQ ID NO 231; 267pp; English.

PT

PT

XX PS

CC

The present invention describes a peptide (I) that is capable of modulating nerve growth factor (NGF) activity. Also described: (1) modified peptide (II) comprising (I) and a vehicle, where the modified peptide is capable of modulating NGF activity; (2) dimer or multimer of (I); (3) modified peptide (III), its multimers or its salt, where the peptide is capable of modulating NGF activity; (4) polynucleotide (IV) encoding (I), (II) or (III); (5) expression vector (V) comprising (IV); (6) host cell (VI) comprising (V); (7) a composition (VII) of matter and a vehicle, where the composition of matter is capable of modulating NGF activity; and (8) pharmaceutical composition comprising (I), (II) or (III) and a diluent or carrier. (I) has analgesic activity, and can be used as an inhibitor of NGF. (I) is useful for treating or preventing a disease or disorder associated with NGF activity by administering (I) to human or animal. The disease or disorder chosen from neurologic pain, painful diabetic neuropathy, post-herpetic neuralgia, inflammatory pain, migraine, asthma, hyperactive bladder, psoriasis, cancer, acute pain, dental pain, pain from trauma, surgical pain, pain resulting from amputation or abscess, causalgia, demyelinating diseases, trigeminal neuralgia, chronic alcoholism, stroke, thalamic pain syndrome, diabetes, acquired immuno deficiency syndrome (AIDS), toxins and chemotherapy, general headache, cluster headache, mixed-vascular and non-vascular syndromes, tension headache, general inflammation, arthritis, rheumatic diseases, lupus, osteoarthritis, inflammatory bowel disorders, inflammatory eye disorders, inflammatory or unstable bladder disorders, skin complaints with inflammatory components, sunburn, carditis, dermatitis, myositis, neuritis, collagen vascular diseases, chronic

```
inflammatory conditions, inflammatory pain associated hyperalgesia and
CC
     allodynia, neuropathic pain and associated hyperalgesia and allodynia,
CC
     diabetic neuropathy pain, sympathetically maintained pain,
     deafferentation syndromes, epithelial tissue damage or dysfunction,
CC
CC
     herpes simplex, post-herpetic neuralgia, disturbances of visceral
CC
     motility at respiratory, genitourinary, gastrointestinal or vascular
CC
     regions, wounds, burns, allergic skin reactions, pruritus, vitiligo,
     general gastrointestinal disorders, colitis, gastric ulceration, duodenal
CC
CC
     ulcers, vasomotor or allergic rhinitis, or bronchial disorders. (I) is
CC
     also useful for modulating pain or promoting analgesia by administering
CC
     (I) to human or animal. (I) is also useful in the manufacture of
CC
     medicament for the treatment of disease or disorder. The present sequence
CC
     is used in the exemplification of the present invention.
XX
SQ
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  Query Match
                          100.0%; Score 104; DB 8; Length 18;
  Best Local Similarity
                          100.0%; Pred. No. 2e-10;
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                                                                 0;
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                                                                     Gaps
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DT
     26-MAR-2002 (first entry)
XX
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     Drosophila melanogaster polypeptide SEQ ID NO 16113.
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KW
     Drosophila; developmental biology; cell signalling; insecticide;
KW
     pharmaceutical.
XX
os
     Drosophila melanogaster.
XX
PN
     WO200171042-A2.
XX
PD
     27-SEP-2001.
XX
PF
     23-MAR-2001; 2001WO-US009231.
XX
PR
     23-MAR-2000; 2000US-0191637P.
PR
     11-JUL-2000; 2000US-00614150.
XX
PA
     (PEKE ) PE CORP NY.
XX
ΡI
     Venter JC, Adams M, Li PWD,
                                  Myers EW;
XX
DR
    WPI; 2001-656860/75.
DR
    N-PSDB; ABL07210.
XX
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CC

PT New isolated nucleic acid detection reagent for detecting 1000 or more genes from Drosophila and for elucidating cell signaling and cell-cell PTPTinteractions. XX PS Disclosure; SEQ ID NO 16113; 21pp + Sequence Listing; English. XX The invention relates to an isolated nucleic acid detection reagent CC capable of detecting 1000 or more genes from Drosophila. The invention is CC useful in developmental biology and in elucidating cell signalling and CC CC cell-cell interactions in higher eukaryotes for the development of CC insecticides, therapeutics and pharmaceutical drugs. The invention discloses genomic DNA sequences (ABL16176-ABL30511), expressed DNA CC CC sequences (ABL01840-ABL16175) and the encoded proteins (ABB57737-CC ABB72072). The sequence data for this patent did not form part of the CC printed specification, but was obtained in electronic format directly CC from WIPO at ftp.wipo.int/pub/published pct sequences XX SQ Sequence 518 AA; Query Match Score 50; DB 4; Length 518; 48.1%; Best Local Similarity 44.4%; Pred. No. 13; 5; Mismatches 8; Conservative 5; Indels 0; Gaps 0; Qу 1 DQLGDWMLNYFRLVPPGT 18 :|: ||| :| :: ||| Db 29 EQIERWMLEWFMVLRPGT 46

Search completed: January 7, 2005, 16:08:12

Job time : 156 secs